



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

NOV 26 1997

Food and Drug Administration  
9200 Corporate Boulevard  
Rockville MD 20850

Mr. Lonnie Witham  
President  
OSMA  
P.O. Box 1846  
Warsaw, Indiana 46581-1846

Dear Mr. Witham:

As you are aware, we have received two citizen petitions (docket numbers 97P-0315/CP 1 and 97P-0315/CP 2). These petitions request that the agency maintain its "policy" of requiring at least two years worth of clinical data prior to filing of a premarket approval (PMA) application for spinal implant devices and that approval be refused for any applications without a substantial amount of this type of data.

A general discussion of the minimum acceptable length of patient follow-up for marketing approval of spinal implant devices will be held on December 11, 1997 at the next Orthopaedic and Rehabilitation Devices Advisory Panel (the Panel) meeting. Interested members of your organization may wish to comment during the scheduled open public session of this meeting. Due to time constraints, the number of commentators will not be large and the time allowed to speak will be kept short. If necessary, this and similar topics may be discussed in more depth at a future Panel meeting.

Interested members should be instructed to contact Ms. Jodi Nashman at 301-594-2036 before December 3, 1997, to request time to speak during the open public session at the next Panel meeting.

If you have any other questions or concerns, please feel free to contact Mr. Mark Melkerson, Branch Chief of the Orthopaedic Devices Branch, at 301-594-2036.

Sincerely yours,

Celia M. Witten, Ph.D., M.D.  
Director  
Division of General and  
Restorative Devices  
Center for Devices and  
Radiological Health

cc: Orthopedic Manufacturers